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(54) Generation of specific probes for target nucleotide sequences

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#### Description

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This invention relates to a method for generating DNA probes specific for an organism and capable of distinguishing between genera and species in a non-empirical manner.

There are many situations where the identification of an organism in a sample is important. This is true in the analysis of clinical, veterinary, food and environmental samples. Currently it is possible to carry out such identification either by classical microbiological methods (where growth characteristics and/or biochemical parameters are monitored), by immunodiagnostic methods or by DNA-based methods.

DNA probes have been used for the identification of organisms as described for example in WO 89/06704 and U. S. Patent No. 4,851,330. Many such probes derive from the observation (see Woese, Scientific American 244 (6) 1981 for review) that parts of the 16S and 23S ribosomal RNA (rRNA) sequences vary in different species. This information was used initially for phylogenetic analyses but it has more recently been used for DNA probe-based methods for the identification of organisms. The method by which the rRNA sequences which are characteristic of an organism were obtained depended initially on differential hybridization experiments in which the target DNA gave a positive signal and the organism from which it was required to be distinguished gave a negative signal or by procedures using hybridization experiments carried out in liquid in which sequences common to be both species are eliminated and sequences (which may be anonymous) which are unique to the organism of interest are retained. A final category of target sequences for DNA probes are regions of the genome which code for some antigen or biochemical product characteristic of the organism.

In all cases the success of the DNA probe depends on its ability to detect a target sequence in the organism of interest while it fails to hybridize to a panel of other organisms that are either closely related to the organism of interest or are likely to occur in the sample under study. Hybridization of a probe to target DNA depends on the DNA sequence and on the hybridization conditions used. There are well established guidelines for the selection of conditions which will allow DNA probes to distinguish between two very closely related sequences (Maniatis, T., et al. (1982) Cold Spring Harbor Publication). The design of DNA probes can be optimized if the DNA sequences targetted have maximum differences from those of other organisms and if a comprehensive data bank of DNA sequences in the region under study is available.

The DNA sequence of a segment of the genome of an organism can be obtained by isolating the DNA segment using a variety of techniques which are widely used by those that employ recombinant DNA methods (see Maniatis, T., et al. supra. More recently methods of amplification of the region of interest using methods such as the Polymerase Chain Reaction (PCR) (Saiki et al. (1985) Science 230 1350-1354) have been described.

The PCR technique requires two oligonucleotide primers that flank the DNA segment to be amplified. Repeated cycles of heat denaturation of the DNA, annealing of the primers to their complementary DNA sequences at a lower temperature, followed by extension of the annealed primer with a DNA polymerase, which may be thermostable, in the presence of the four deoxyribonucleotides gives rise to specific DNA sequences in sufficient quantities to be manipulated further.

In one use of the PCR method Chen, K., et al. (FEMS Microbiology Letters (1989) 57, 19-24) mplified E. coli using primers derived from the regions of the 16S rRNA gene which tend to be conserved in a variety of organisms examined. A similar approach has been used by Medlin, L., et al. Gene (1988) 71, 491-499) who amplified eucaryotic rRNA coding regions for a phylogenetic study.

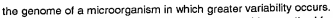
WO 88/03957 describes a method of constructing probes for the detection of intragenic rRNA sequences to distinguish non-viral organisms. The probes are generated from the 16S or 235 rRNA by making a cDNA copy of the rRNA using reverse transcriptase according to the method of Lane, D.J. et al. (Proc. Natl. Acad. Sci. U.S.A. (1985); 82, 6955-6959) and thereafter determining the sequence of the cDNA by standard methods.

Göbel, U.B. et al. (Journal of General Microbiology (1987); 133, 1969-1974) also disclose the construction of species-specific probes to variable regions within the 16S-like and 235-like rRNA genes using synthetic oligonucle-

In all of the references cited above the variable regions are intragenic.

The choice of a target sequence for a probe currently involves a) the identification of an area of sufficient interspecies diversity or variation that will allow for the provision of a specific probe and b) a target which is preferably present in the organism in a high number of copies. The rRNA gene products (16S and 23S) appear to fulfil both of these criteria and as such have been the target for many studies and, indeed, DNA probe kits directed to those regions are available commercially for some organisms. Most comparisons to date have been between the rRNA genes from different genera and these have highlighted a pattern of variable regions within the gene flanked by adjacent more conserved regions.

We have found that when related species are compared the "variable" regions are occasionally very similar (see Example 2), if not identical. This highlights the need for a method to obtain sequence data from a catalogue of organisms to allow one to select the correct probe sequence and the appropriate hybridization conditions or to identify regions of



It is an object of the present invention to provide a method for obtaining DNA sequences which can be used to provide a DNA data base for the choice of probe and hybridization conditions in a rapid and useful manner.

It is a further object of the present invention to identify new target areas that contain a high degree of diversity between organisms and to generate highly specific DNA probes thereto in a variety of organisms of interest to the clinician, veterinary practitioner, the food technologist and the environmentalist.

Accordingly, the invention provides a method for generating a DNA probe specific for an organism to be identified and capable of distinguishing between genera and species in a non-empirical manner, said method comprising the steps of:

- a) amplifying, using a pair of oligonucleotide primers, a variable intergenic region of the genome present in the organism to be identified and in a number of organisms phylogenetically related to, or suspected of being present in a given sample containing said organism to be identified and having said variable region in its genome, said primers corresponding to regions or portions thereof flanking said intergenic region and known or suspected of being conserved in said organisms;
- b) determining the sequence of the amplified region;

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- c) selecting said sequence or a portion thereof to generate said probe specific for said organism to be identified by comparison with said other amplified regions; and
- d) defining the hybridization conditions required to obtain a specific signal based on the precise nucleotide sequence of the selected probe.

The method in accordance with the invention enables one to generate DNA probes which can discriminate between very closely related organisms. Thus the method according to the invention can distinguish between organisms in which only 2 bases are different at a given gene locus as hereinafter described.

The method according to the invention has universal application in the generation of DNA probes to variable regions of the genome of a given organism. The oligonucleotide primers may correspond to DNA sequences of conserved regions or portions thereof adjacent said variable region or portion thereof.

The method in accordance with the invention can be used to target an unknown region of the genome by rapidly establishing a DNA sequence data bank for that region for a number of organisms which are likely to be present in a sample and by deriving a DNA probe fragment which will identify the organism of interest under hybridization conditions determined by the sequence of the probe and the extent to which it differs from the other organisms. The method according to the invention represents a significant improvement over existing methods for identifying sequences useful as DNA probes.

Because of the inadequacies in some cases of the 16S rRNA gene as a locus for DNA probing we have availed of the universality of the method in accordance with the invention to analyse other genomic regions for their ability to provide informative targets for DNA probes. We have searched in the first instance for regions that are least likely to be conserved. This consideration led us to focus attention on those regions of the genome which are intergenic, viz. spacers between functional genes as there should be minimal evolutionary pressure to conserve there regions.

Previous authors have published data on the intergenic region between the 16S and the 23S rRNA for <u>E. coli</u> (Brosius <u>et al.</u> J. Mol. Biol. <u>148</u> 107-127 (1981)), <u>Streptomyces lividans</u> and <u>Mycobacterium bovis</u> BCG (Suzuki <u>et al.</u> J. Bact. 170 2886-2889 (1988)) and <u>Bacillus subtilis</u> (Green <u>et al.</u> Gene <u>37</u>, 261-266 (1985)). Examination of these sequences by us has shown that there is a great variation between these unrelated genera.

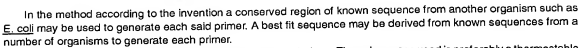
According to the invention the variable region is a variable intergenic region and the primers correspond to conserved regions or portions thereof flanking said intergenic region. Thus as an example one of the primers may correspond to a conserved region of the 16S rRNA gene and the other primer may correspond to a conserved region of the 23S rRNA gene.

Alternatively, one of the primers may correspond to the conserved 3' end of the 16S rRNA gene and the other primer may correspond to the conserved 5' end of the 23S rRNA gene.

The primers can be derived from the available sequence data for 16S and 23S genes. When such primers are amplified in accordance with the invention using the PCR technique the amplified (intergenic) region can, in some situations, vary in size, such is the diversity between species, as hereinafter described.

By DNA sequencing such amplified regions additional detailed information on panels of microganisms has been obtained which has allowed DNA probes to be prepared which can distinguish between closely related organisms.

Although the intergenic region between the 16S and 23S genes is the target used in Example 1 other intergenic regions which are bounded by known sequences of functional genes can also be used as targets for DNA probes in accordance with the invention.



The amplification is preferably carried out by the PCR technique. The polymerase used is preferably a thermostable polymerase such as <a href="https://documents.org/nc/">Thermus aquaticus</a> (Taq) enzyme. It will be appreciated that zones of the genome of an organism which have been identified as target sequences for DNA probes in accordance with the invention may be isolated using methods other than PCR amplification.

Following amplification and prior to sequencing, the amplified nucleotide sequence may be ligated to a suitable vector followed by transformation of a suitable host organism with said vector. One thereby ensures a more readily available supply of the amplified sequence.

Alternatively, following amplification, the amplified sequence or a portion thereof may be chemically synthesized for use as a nucleotide probe. In either situation the DNA sequence of the variable region is established using methods such as the dideoxy method (Sanger, F. et al. Proc. Natl. Acad. Sci (1977) 74, 5463-5467). The sequence obtained is used to guide the choice of the probe for the organism and the most appropriate sequence(s) is/are selected.

We have determined that the DNA used for amplification may come from autoclaved samples. This is a particularly useful demonstration as it allows for ready preservation of samples and greater safety for laboratory workers.

The target organism is suitably a microorganism. For example, the organism may be selected from a species of Aeromonas, Bacillus, Clostridium, Enterococcus. Eschericia, Klebsiella, Mycobacterium, Pseudomonas, Salmonella Serratia, Staphylococcus or Streptococcus. Specific microbial species within these species include Aeromonas caviae, Aeromonas hydrophila, Aeromonas media, Aeromonas salmonicida, Aeromonas sobria, Bacillus subtilis, Clostridium butyricum, Clostridium difficile, Clostridium pasteurianum, Clostridium perfringens, Clostridium tyrobutyricum, Eschericia coli, Enterococcus faecalis, Klebsiella pneumoniae, Mycobacterium avium, Mycobacterium bovis, Mycobacterium tuberculosis, Pseudomonas aeroginosa, Pseudomonas fluorescens, Salmonella typhimurium, Serratia mercescans, Staphylococcus aureus and Streptococcus pyogenes.

The invention also provides specific probes obtainable by the preceding method.

The invention provides a variety of probes for variable intergenic regions intermediate the genes coding for 16S rRNA and 23S rRNA and derived from the following sequences or sequences complementary thereto:

#### Clostridium difficile -

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#### Clostridium pasteurianum -

#### Clostridium perfringens -

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#### Mycobacterium avium -

#### Mycobacterium bovis -

ACCGGAAAGGTCCGTGCGTGAAATTTAACCTTCCTTCCCTTTTCTAAGGAGCACCACGA
AAACGCCCCAACTGGTGGGCGTAGGCGTGAGGGGTTCTTGTCTGTAGTGGGCGAGACGG
GGTGCATGACAACAAAGTTGCCACCAACACACTGTTGGGTCCTGAGGCAACACTCGGAC
TTGTTCCAGGTGTTGTCCCCACCGCCTTGGTTGGTGGGGTGTGTTTTGAGAACTGG
ATAGTGGTTGCGAGCATCAATGGATACGCTGCCGGCTAGCGGTGGCGTGTTCTTTGTGC
AATATTCTTTGGTTTTTGTTGTGTTTTGTAAGTGTCTAAAGGGCGCAT

#### Mycobacterium tuberculosis -

ACCGGAAGTCGTCGGGATCACCTCCTTTCTAAGGAGCACCACGAAAACGCCCCAACTGG
TGGGTCAGGCGTGAGGGGTTCTTGTCTGTAGTGGGCGAGACGGGGTGCATGACAACAAA
GTTGGCCACCAACACACTGTTGGATCCTGAGGCAACACTCGGACTTGTTCCAGGTGTTG
TCCCACCGCCTTGGTGGTGGTGGTGGTGTTTTGAGAACGTGATAGTGGTTGCGAGCATCA
ATGGATACCCGTGCCGGCTAGCGGTGGCGTGTTCTTTGTGCAATATCTTTGGTTTTTGT
TGTGTTTGTAAGTGTCTAAGGGCGCA

The above specifically mentioned probes are just some examples of specific probes which can be generated in accordance with the method of the invention.

To help satisfy the second requirement of DNA probes (i.e. high copy number) non-translated, transcribed regions have been chosen for analysis because these fulfil a functional role other than that of a messenger RNA which must be translated. Because of the usefulness of the intragenic regions of ribosomal genes other intragenic regions were

investigated as being appropriate targets for DNA probes.

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To allow us to define probes which can distinguish between related genera and species we have used the integrated method of the invention to rapidly define DNA probes for closely related organisms (see comparative Example 2). In one demonstration of the method in which the intragenic region of the 16S rRNA was targeted we define a probe/ hybridization conditions couple in which sequences differing only by 2 bases are distinguished.

To demonstrate the general usefulness of the method we have used available DNA sequences from <u>E. coli</u> to provide primers for a wide range of Gram positive, Gram negative, aerobic and anaerobic organisms.

Although the variable regions of the gene coding for 16S rRNA have been the basis of species-specific probes, the use of the constant regions as primers for PCR allows one to rapidly isolate that part of the gene coding for 16S rRNA from species on which no sequence or hybridization analysis has been performed. The subsequent DNA sequencing of the amplified region yields information on the variable part of the organism under study and this can be used as a DNA probe in subsequent experiments under defined conditions.

The above specifically mentioned probes are some further examples of specific probes which can be generated in accordance with the method of the invention.

The data provided hereafter in Example 2 and above were based on sequences in the intragenic region of the 16S rRNA gene. Other RNA molecules that have a functional role other than that of a template for translation should also be useful as target sites for the generation of specific probes against RNAs which are present in high copy numbers. An example of such a transcribed but optionally translated gene region is the ribonuclease P component RNA (M1). A study of the published data (Bryan, D. James et al. (1988) Cell 52, 19-26) showed that there are variable regions within this RNA. We have demonstrated (Example 3) that effective primers can be prepared from the available data that will allow amplification of these variable regions. Using the methodology according to the invention one can derive a panel of DNA sequences of the amplified region and, using the method herein specified in accordance with the invention, develop DNA probes based on a comparison of the DNA sequences obtained.

The present invention enables the intergenic variable regions to be analysed and to provide a DNA probe which will hybridize specifically under selected conditions to the chosen DNA.

Inherent in the method according to the invention is the selection of hybridization conditions appropriate for any given probe in a non-empirical manner. The hybridization conditions are selected on the basis of the precise nucleotide sequence of the probe which can be determined in a manner known per se for stable hybridization, so as to distinguish the sequence selected from that of closely related species as hereinafter further described in the Examples. The biochemical and physical parameters selected for optimal hybridization in any given situation are based on the ratio of A: T to G.C pairings involved in hybridization and modifying principally the temperature and salt concentration of the hybridization medium accordingly. Determination of the optimal hybridization conditions involves comparison of the amplified sequence with closely related sequences in an available or freshly created data bank, the latter having been created by the determination of previously unknown sequences in accordance with the invention. Thus the method according to the invention has both increased sensitivity due to the determination of optimal and controlled hybridization conditions and specificity due to the generation of species-specific probes appropriate to any region of the genome of an organism.

The method according to the invention provides for the generation of DNA probes for the detection of a variety of species of organisms and probes which can detect many members of given genera. The present invention provides a more general demonstration of target sites in organisms and thus represents an improvement on current targets for DNA probes and a logical and data based selection of DNA probe sequences and the conditions of hybridization under which they should be used.

The probes obtained in accordance with the method of the invention may be used for the detection and/or determination of an organism containing or suspected of containing a target nucleotide sequence by contacting said organism with said probe having a sequence complementary to said target sequence.

The applications of the above probes will be of benefit in a variety of situations. The probe for Mycobacterium bovis, for example, will provide a non-subjective test for the presence of this organism in cattle (or other species). Currently bovine T.B. is a disease of major economic importance in some European countries and in very many developing countries. The description of a probe for Mycobacterium tuberculosis will allow the detection of this fastidious organism in humans (currently it is undetected by standard microbiological procedures in 50% of clinically positive cases) and this is of growing importance as it occurs in about 20% of persons suffering from full-blown AIDS.

A probe for Aeromonas salmonicida will allow the detection in the environment or in fish of this pathogen which causes furunculosis with major economic losses in the salmon farming industry. Finally, as further examples inter alia of the usefulness of these probes, the sequences provided for various clostridia will be of benefit in clinical analyses (e.g. Clostridium difficile) or the food industry.

The invention further provides a method of detecting a specific organism in a mixture of organisms, which method comprises determining the sequences of two selected nucleotide regions of said specific organism by a sequence of steps set forth in the method hereinbefore specified, and using said sequenced regions to generate a pair of primers

to selectively amplify a specific nucleotide region of said organism and thereby detect said specific organism. The organism is suitably any one of those specifically hereinbefore mentioned.

It will be appreciated that the above method may comprise determining the sequences of two selected variable regions and using said sequenced variable regions to generate a pair of primers to selectively amplify a specific variable region and, thereby, generate a species-specific probe for said specific organism or, alternatively, to detect the product of amplification. When two variable regions are obtained they can serve as very specific primers for the particular organism of interest. It will be appreciated the primers thus produced are equivalent to probes as hereinbefore described.

Alternatively, a combination of one species-specific primer as hereinbefore described and a constant or conserved region primer can be used in combination to specifically amplify the species of interest.

The selective amplification of the species of interest in the manner described above can further be confirmed by the use of a probe specific for the species.

It will be appreciated that the methods according to the invention may be 'one-tube' methods involving minimal equipment and fewer manipulations than heretofore. Accordingly, the invention further provides any method hereinbefore specified wherein DNA preparation and amplification are carried out in a single tube.

The methods in accordance with the invention have application in animal, plant and microbial organisms. It will be appreciated that the probes prepared in accordance with the invention can be used as biosensors for a variety of applications.

In the accompanying drawings:

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Fig. 1 is a photograph of an agarose gel amplified 16S-23S intergenic regions for <u>Clostridium</u> species prepared in Example 1;

Fig. 2 is a photograph of an autoradiogram of hybridization of a DNA probe for the 16S-23S intergenic region of Mycobacterium bovis to Mycobacterium bovis and Mycobacterium avium DNA;

Fig. 3 is a photograph of an agarose gel of amplified 16S rRNA regions from a panel of organisms as prepared in comparative Example 2;

Fig. 4 is a photograph of an autoradiogram of hybridization of a DNA probe for 16S rRNA intragenic regions of Aeromonas salmonicida to a panel of organisms; and

Fig. 5 is a photograph of an agarose gel of amplified ribonuclease P regions from a panel of organisms as prepared in comparative Example 3.

The invention will be further illustrated by the following Examples.

#### **EXAMPLE 1**

General Procedure for the Isolation and PCR Amplification of DNA from Microorganisms: Illustration of Method applied to Intergenic Regions

50 μl (105-106 bacteria) of a fresh culture were pelleted by centrifugation at 10,000 r.p.m. for 1 min. in a bench top centrifuge and the supernatant discarded. The bacterial cultures were obtained from various laboratories in the Department of Microbiology, University College Galway, Ireland, University College Hospital, Galway, Ireland and the Veterinary College, University College Dublin, Ireland. The pellet was then resuspended in 25 μl of water. The sample was heated to 95°C for 10 min. in a 750 μl Eppendorf tube, centrifuged briefly to remove condensation from the lid of the tube and then incubated in the presence of proteinase K (at a concentration of 50 μg/ml) at 55°C for 15 min. Proteinase K was then denatured at 95°C for 15 min. Condensation was again removed from the Eppendorf lid by a brief centrifugation and DNase free RNase A (Sigma) was added to a final concentration of 10 μg/ml and allowed to incubate for 15 min. at 37°C to denature RNA. These incubations and reactions were carried out using a Perkin-Elmer-Cetus Thermo-cycler.

After RNA digestion, a 2 x PCR reaction buffer (cocktail) (100mm KCI, 20mm Tris, pH 8.3, 3mm MgCl<sub>2</sub>, 0.2% gelatin, 400 μm dNTP, an appropriate concentration of primers (~500 pmol of each for a 20 mer), and 2.5 units of Taq polymerase were added and 30 PCR cycles, in a final volume of 50 μl, were then carried out. Typically, a PCR cycle consists of DNA heat denaturation at 94°C for 1 min., annealing of primers at 37°C-55°C for 2 min. and an extension reaction period at 72°C for 1-3 min. The samples were allowed to cool to room temperature gradually and 1/10 (5 μl) of the reaction were analysed by gel electrophoresis on a 4% Nu-Sieve (Nu-Sieve is a Trade Mark) agarose mini-gel

to determine the success of amplification (see Figs. 1, 3 and 5).

Alternative methods using a combination of proteinase K, detergents, phenol extraction and ethanol precipitation can also be used successfully for the preparation of the DNA.

20-Mer oligonucleotide primers which come from the 3' end of the 16S rRNA gene and the 5' end of the 23S rRNA gene were synthesized on an Applied Biosystems or Beckman (Applied Biosystems and Beckman are Trade Marks) oligonucleotide synthesizer. The primers thus synthesized correspond to conserved sequences in those abovementioned regions as defined by Clustal Analysis (Higgins et al., Gene, 73 237-244 1988) of available 16S and 23S rRNA

The sequence of the 16S rRNA 3' primer is

AGTCGTAACAAGGTAGCCGT

The sequence of the 23S rRNA 5' primer is:

T/G A/G C/T TGCCAA G/C GCATCCACC

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where "/" indicates a degenerate selection of a base at that position.

After deblocking, purification of the oligonucleotides was carried out by elution from preparative polyacrylamide

gels, and purified further by chromatography on NAP10 columns.

PCR amplification with the above primers was carried out using the conditions outlined above for the following microorganisms: Aeromonas caviae, Aeromonas hydrophila, Aeromonas salmonicida, Aeromonas sobria, Bacillus subtilis, Clostridium butyricum, Clostridium difficile, Clostridium pasteurianum, Clostridium perfringens, Clostridium tyrobutyricum, Escherichia coli, Enterococcus faecalis, Mycobacterium avium, Mycobacterium bovis, Mycobacterium tuberculosis, Pseudomonas aeroginosa, Serratia mercescans, Staphylococcus aureus and Streptococcus pyogenes. The sizes of the amplified regions ranged from 200 to 650 base pairs. The data for the five Clostridium species are shown in Fig. 1 in which Lanes 1-6 represent the following:

Clostridium perfringens amplified 16S-23S rRNA intergenic regions Lane 1:

Clostridium difficile amplified 16S-23S rRNA intergenic regions Lane 2:

Clostridium pasteurianum amplified 16S-23S rRNA intergenic regions Lane 3:

Clostridium butyricum amplified 16S-23S rRNA intergenic regions Lane 4:

Clostridium tyrobutyricum amplified 16S-23S rRNA intergenic regions Lane 5:

pBR Hae III size markers Lane 6:

The PCR amplified DNA was subcloned into the vector M13 mp II by blunt end ligation into the Smal site. The inserted DNA was sequenced by the Sanger dideoxy chain termination method. These operations were carried out by subjecting 50 ng of amplified DNA (approx. 20 µl of the PCR reaction) to drop dialysis against a 1000-fold volume of TE buffer (10 mM Tris, pH 8, 1 mM EDTA) for 2 h. The DNA was dried in a Heto-Vac (Heto-Vac is a Trade Mark) dessicator and then resuspended in 10 µl of ligation buffer containing 10 ng of Smal digested M13 Mp 11 and one unit of T4 ligase. Ligation took place at 10°C for at least 5 h. The DNA mixture was used to transform E. coli JM201. A single stranded DNA was prepared from recombinants for dideoxy sequencing using T7 polymerase. The DNA sequences that were obtained are set out above.

A clustal alignment of the Mycobacterium species sequences is shown in Table 1. The variable regions between them are indicated by a lower density of conserved bases (asterisk). Many possibilities for probe generation arise from this analysis. One fragment, 20b (which encompasses a 9b difference) underlined was selected (in Table 1) to demonstrate that Mycobacterium bovis can be distinguished from Mycobacterium avium (Mycobacterium avium is used as an indicator of non-pathogenic infections in cattle in current tests for bovine T.B.). When this sequence was radioactively labled by polynucleotide kinase (Amersham) in the presence of  $\gamma^{32}$ PATP and used as a probe to the amplified intergenic regions of Mycobacterium bovis and Mycobacterium avium that had been slot blotted to a Nytran (Nytran is a Trade Mark) membrane (Schleiser and Schuell ..), the Mycobacterium bovis was detected whereas the Mycobacterium avium was not (Fig. 2). in Fig. 2 "A" corresponds to Mycobacterium bovis and "B" corresponds to Mycobacterium avium. The conditions for hybridization were 55°C for 2 h. in 6 x SSPE, 0.1% S.D.S. and washes were twice at room temperature in 6 x SSC, 0.1% S.D.S. and once for 2 min. at 55°C in 6 x SSC 0.1% S.D.S. DNA sequences for Clostridium species

obtained in a similar manner are shown in Table 2. Again the possibilities for DNA probes are obvious from the clustal analysis. An analysis of the sequences shown in Table 2 for three species of <u>Clostridium</u> show regions in which an identical series of sequences (indicated by asterisks) occurs which could serve as the basis of a genus specific probe for Clostridium, if required.

#### **EXAMPLE 2** (Comparative)

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## Amplification of V2 and V6 regions of microorganisms and probe generation therefrom.

Amplification was carried out in accordance with the procedures set out in Example 1. The 20mer oligonucleotide primers which flank each side of the V2 (R1/R2) and V6 (U1/U2) variable regions of the 16S rRNA gene were selected from available data [Dams et al. Nucleic Acids Research Supplement 16 r87-r173 (1988)].

The primers thus synthesized had the following sequences:

15	5'	3'
	R1: AATTGAAGAGTTTGATCATG	3'
20	R2: ACATTACTCACCCGTCCGCC	
	5'	3 '
25	U1: GCAACGGCGAAGAACCTTAC	
		31
30	5'	,
	U2: GACAGCCATGCAGCACCTGT	

PCR amplification with R1/R2 and U1/U2 as primers for V2 and V6, respectively, was carried out using the conditions outlined above for the following microorganisms: <u>Aeromonas salmonicida</u>, <u>Clostridium difficile</u>, <u>Klebsiella pneumoniae</u>, <u>Mycobacterium bovis</u>, <u>Pseudomonas fluorescens</u>, <u>Salmonella typhimurium</u> and <u>Staphylococcus aureus</u>. R1/R2 amplification of these species gave rise to approximately 120 base pair fragments in all cases analysed. U1/U2 primer amplification gave rise to an approximately 100 base pair fragment. Fig. 3 depicts the results obtained for R1/R2 primer V2 amplification and U1/U2 primer V6 amplification in which the lanes 1-16 represent the following:

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	Lane 1:	pBR Hae III size markers.
	Lane 2:	Aeromonas salmonicida V2 amplification.
	Lane 3:	Aeromonas salmonicida V6 amplification.
	Lane 4:	Clostridium difficile V2 amplification.
45	Lane 5:	Clostridium difficile V6 amplification.
	Lane 6:	Klebsiella pneumoniae V2 amplification.
	Lane 7:	Kmebsiella pneumoniae V6 amplification.
	Lane 8:	Mycobacterium bovis V2 amplification.
	Lane 9:	Mycobacterium bovis V6 amplification.
5 <b>0</b>	Lane 10:	Pseudomonas fluorescers V2 amplification.
	Lane 11:	Pseudomonas fluorescers V6 amplification.
	Lane 12:	Salmonella typhimurium V2 amplification.
	Lane 13:	Salmonella typhimurium V6 amplification.
	Lane 14:	Staphylococcus aureus V2 amplification.
55	Lane 15:	Staphylococcus aureus V6 amplification.
	Lane 16:	pBR Hae III size markers.

The amplified bands indicate that the constant region primers are effective for Gram positive, Gram negative,

aerobic and anaerobic organisms.

DNA sequences were obtained as described in Example 1. Examples of some of the DNA sequences obtained for both the V2 and V6 region are shown in Tables 3a and 3b. The sequences for the organisms other than <u>E. coli</u> were obtained using the methods herein described in accordance with the invention. The regions which correspond to the universal primers are boxed. The variable regions between them are indicated by a lower density of conserved bases (asterisk). A DNA probe specific for <u>Aeromonas salmonicida</u> selected on the basis of this analysis is indicated by a dashed line (Table 3b). This V6 probe sequence successfully distinguished <u>Aeromonas salmonicida</u> from <u>E. coli, Salmonella typhimurium, Clostridium difficile, Aeromonas hydrophila, Aeromonas media and <u>Aeromonas caviae</u>, when these cultures were amplified using primers for the constant regions flanking V6 (see Fig. 4). It will be noted that the hybridization conditions used (hybridization was carried out at 57°C for 2 h. in 6 x SSPE, 0.1% SDS with the most stringent wash at 60°C for 30 min. in 2 x SSC) permitted the probe to distinguish between <u>A. salmonicida</u> and <u>A. hydrophila</u>, in which there is only a 2-base pair difference.</u>

In Fig. 4 Lanes 1-7 represent the following:

15 Lane 1: Aeromonas salmonicida

Lane 2: Aeromonas hydrophila

Lane 3: <u>Aeromonas media</u>

Lane 4: Aeromonas caviae

Lane 5: <u>Escherichia coli</u>

20 Lane 6: Salmonella typhimurium

Lane 7: Clostridium difficile

#### **EXAMPLE 3** (Comparative)

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### Amplification of Variable Intragenic Regions of Ribonuclease P RNA Gene

Using the methods described in Examples 1 and 2, DNA primers selected as conserved from an analysis of published data (Bryan, D. James et al. (1988) supra) which flank the variable intragenic regions of ribonuclease P were synthesized. The sequences of the primers are:

synthesized. The sequences of the primers are:

A 17-mer oligonucleotide

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3'

G A/T C/T C A/G G A/G C/T A A/G TCGC T/C GC

and
a 19-mer oligonucleotide

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י 3

## C/T C G/T ATAAGCC G/A G/T GTT T/C TGT

These were used to amplify the corresponding intragenic regions in <u>Aeromonas salmonicida</u>, <u>Clostridium difficile</u>, <u>Clostridium pasteurianum</u>, <u>Escherchia coli</u>, <u>Enterococcus faecalis</u> and <u>Serratia mercescans</u>. The result (shown in Fig. 5) shows that both Gram positive, Gram negative, anaerobic and aerobic organisms can be amplified using these primers. In Fig. 5 Lanes b-h represent the following:

Lane b: Escherchia coli

Lane c: Enterococcus faecalis

Lane d: <u>Serratia mercescans</u>

Lane e: <u>Aeromonas salmonicida</u>

Lane f: Clostridium pasteurianum

Lane g: Clostridium difficile

Lane h: Ikb ladder size marker.

The arrow indicates Ribonuclease P amplified DNA.

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## TABLE 1

Mycobacterium tuberculosis	Mycobacterium tuberculosis	Mycobacterium tuberculosis	Mycobacterium tuberculosis	Mycobacterium tuberculosis
Mycobacterium bovis	Mycobacterium bovis	Mycobacterium bovis	Mycobacterium bovis	Mycobacterium boxis
Mycobacterium avium	Mycobacterium avium	Mycobacterium avium	Mycobacterium avium	Mycobacterium avium
TTTTGTTGTTGTTAAGTGTCTAA-GGGCGCA- TTTTGTTGTTGTTAAGTGTCTAAAGGGCGCAT TCTTGT-GTGAAGTAAGTGTTTAAGGGGGG-AT	GATAGTGGTTGCGAGCATCAATGGATACCCGTGCCGGCTAGCGGTGGCGTGTTCTTTGTGCAATAT-CTTTGGT GATAGTGGTTGCGAGCATCAATGGATACGC-TGCCGGCTAGCGGTGGCGTGTTCTTTTGTGCAATATTCTTTTTGT GATAGTGGTTGCGATGATCTAGGTGAGCGCATGGTCTTTCGTTGCCCGGCCGTTGATCGAAATGTGTAATTTCTTTTTAAC ************************	GATCCTGAGGCAACACTCGGACTTGTTCCAGGTGTTGTCCC-ACCGCCTTGGT-GGTGGG-TGTG-GTGTTTGAGAACTG GGTCCTGAGGCAACACTCGGACTTGTTCCAGGTGTTGTCCCCCACCGCCTTGGTTGG	-CAGGCGTGAGGGGTTCTTGTCTGTAGTGGGCGAGACGGGGTGCATGACAAAGTTGGCCACCACCACCACTGTTG GTAGGCGTGAGGGGTTCTTGTCTGTAGTGGGCGAGACGGGGTGCATGACAAAAGTTG-CCACCAACAACACTGTTG TCGGGAGCCAGTAGGGGTTCCCGTCTAGTGACGGGGGCCCGGGTGCGCAACAAAAGTAGATTGCCAGACACACTATTG TCGGGAGCCAGTAGGGGTTCCCGTCTAGTGACGGGGGCCCGGGTGCGCAACAAAATGATTGCCAGACACACTATTG	ACCGGAAGTC-GT-CGGGATCACCTCCTTTCTAAGGAGCACCACGAAAACGCCCC-AACTGGTGGGT ACCGGAAAGGTCCGTGCGTGAAATTTAACCTTCCTTCCCTTTCTAAGGAGCACCACGAAAACGACCCCGAACTGGTGGGC ACCGGAGGTGGGGC-TGATCACCTCCTATTCTAAG-AGCACCACAAAACGACCCCGAACTGGTGGGG ******

Clostridium perfringens Clostridium difficile Clostridium pasteurianum

	<u>-</u>		
Clostridium perfringens Clostridium difficile Clostridium pasteurianum	Clostridium perfringens Clostridium difficile Clostridium pasteurianum	Clostridium perfringens Clostridium difficile Clostridium pasteurianum	
TAATATAGGTCAAGCTACAAAGGGCGCAT CTTTGTAAGAATCAATT-TAAGGTCAAGCTACAAAGGGCGCAT TAACGCCTAAAAGGATAACAAGGTCAAGCTACAAAGGGCGCAT ************************************	AAACAACAAGCCAAATTGGCAAAAACCAATTTCTATTCTTTGTAAAATGAGAACTATAACAAATCAAACAAGCCAAACTATAAAAATCAAATTGGTAAAATTTTTGGATGCCTAGCTACGTTCTTTGAAAATTGCACAGTGAATAAAATCAAATAAGGCTAAAACTCTAAAAACGGGCAAAAATAAAATCCTTTGATAAT-ATGATTTTAATCGAAAAAGATTGAAATTAA-ACAATAAAGACTAAAACTCTAAAAACGGGCAAAAAGAATTAAATCCTTTGATAAT-ATGATTTTAATCGAAAAAGATTGAAATTAA-ACAATAAAGACTAAAACTGGGCAAAAAGGACTAAAACGGGCAAAAAGATTAAAAAAGACTAAAAACGGGCAAAAAAGATTAAAAAAGATTAAAAAAGACTAAAAAAAA	AATATCTCTGTTTAATTTTGAGAGACTATCTCTCAAAATTGTTCTTTGAAAATTGCACATAATTTAATT	

TABLE

AGGAGAACTGC-GGCTGGATCACCTCCTTTCTAAGGAATA--CATCTTAGGACAACTAAGATG-ATAATGAATTCTGGATAGAAACCTGCCGG-TGAATCACCTCCTTTCTAAGGAG-AATAGAAAAGAAGAAAATTCTT-TCTAAA--GG---CTGAAT

## тавье за

Multiple alignment of the DNA sequences of the V2 loop (region) of the gene coding for 16S rRNA of the following organisms: Beromonas salmonicida, Clostridium perfringens, Clostridium difficile, Salmonella typhimurium. Klebsiella pneumoniae, Rseudomonas fluorescens and Staphylococcus aureus to the known E. coli V2 sequence.

# Aeromonas salmonicida Escherichia coli

Clostridium perfringens Clostridium difficile Pseudomonas fluorescens Salmonella typhimurium Staphylococcus aureus Klebsiella pneumoniae

 ${ t AATTGAAGAGTTTGATCATG}$ CCTC ${ t GAACGCTGAACGCCTGGCGCAACACACATGCAAGTCGAGCGGCAGCGGGAAAGTA}$ Primer Region R1
AATTGAAGAGTTTGATCATG CCTCAGATTGAACGCTGGCCGGCAGGCCTAACACATG-AAGTCGAACGGTAACAGGAA-GAA

 $\mathtt{AATTGAAGAGTTTGATCATG}$ CCTCAGATTGAACG-TGGCGGCAGGCCTAACACATGCAAGTCGAACGGTAACAGGAA-GCA AATTGAAGAGTTTGATCATG|GCTCAGGATGAACGCTGGCGGCGTGCCTAACACATGCAAGTTGAGCGAT--------aattgaagagtttgatcatg|gctcaggatgaacgctggcggcgagcttaacacatgcaagtcgagcgat------gaa AATTGAAGAGTTTGATCATG|GCTCAGATTGAA--CTGGCGGCAGGCCTAACACATGCAAGTCGAGCGGTA----GAGAGAA AATTGAAGAGTTTGATCATG|GCTCAGATTGAACGCTGGCGGCAGGCCTAACACATGCAAGTCGAGCGGTAC--GACAGAGA **水水水水水水水水水水水水水水水水水水水水水水水** --ttgaagagtttgatcatg|gctcaggatgaactctggcggcgagcctaatacatgcaagtcgagcgaacg--gacgagaa \*\*\* \*\*\*\*\* 存者 计操作 计计算计算 计计算计 计计

<u>Escherichia coli</u> Pseudomonas fluorescens Clostridium perfringens seromonas salmonicida <u>Klebsiella pneumoniae</u> Salmonella lostridium difficile typhimurium

GCTTGCT-TCTTTGCTG-ACGAGT GGCGGACGGGTGAGTAATGT

Primer Region R2

GCTTGCTACTTTTGCCG-GCGAGC GGCGGACGGGTGAGTAATGT

GGCGGACGGGTGAGTAATGT

GGCGGACGGGTGAGTAATGT \*\*\*\* GGCGGACGGGTGAGTAATGT

G-TTTCTTCGGGAAATGGATTAGC

Staphylococcus aureus GCTTGCT-CTCG-GGTG-ACGAGC GGCGGACGGGTGAGTAATGT GCTTGCT-CGTTCGCTG-ACGAGT GGCGGACGGGTGAGTAATGT --ttacttcggtaaa--ga---gc ggcggacgggtgagtaatgt GCTTGCTTCTCTTGA----GAGC GCTTGCTTCTCT-GATG--TTAGC

\*mean base conserved in all cases studied

-indicates a gap which allows for optimal alignment

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## TABLE 3b

Murtiple alignment of the DNA sequences of the V6 loop (region) of the gene coding for 16S rRNA of the following organisms: <u>Beromonas salmonicida</u>, <u>Beromonas hydrophilia</u>, <u>Clostridium perfringens</u>, <u>Clostridium</u> difficile. Salmonella typhimurium, Klebsiella pneumoniae. Pseudomonas fluorescens and Staphylococcus aureus to the known E. coli V6 sequence.

# Aeromonas salmonicida Escherichia coli

GCAACGCGAAGAACCTTACC|TGGCCTTGACATGTCTGGAA-TCCTGTAGAGATA---

Primer Region Ul

GCAACGCGAAGAACCTTACC|TGGCCTTGACATGTCTGGAA-TCCTGCAGAGATG---

-----cgggaatca g aac-a

--cgggaatca|g aac-a

GCAACGCGAAGAACCTTACC|TGGTCTTGACATCCACGGAAGT-TTTCAGAGATGAGAATGTGCCTTCGGGAACCC TGA-GA

Pseudomonas fluorescens Klebsiella pneumoniae <u>Salmonella typhimurium</u> Clostridium difficile Clostridium perfringens Beromonas hydrophilia Staphylococcus aureus

Primer Region U2
CAGGTGCTGCATGGCT CAGGTGCTGCATGGCT CAGGTGCTGCATGGCT

eromonas salmonicida scherichia coli Pseudomonas fluorescens Klebsiella pneumoniae Salmonella typhimurium Clostridium difficile eromonas hydrophilia Staphylococcus aureus ostridium perfringens

CAGGTGCTGCATGGCT CAGGTGCTGCATGGCT CAGGTGCTGCATGGCT CAGGTGGTGCATGGCT CAGGIGCIGCATGGCI CAGGTGATGCATGGCT 计不可处计数 经非非非非非非非

\*mean base conserved in all cases studied

indicates a gap which allows for optimal alignment

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GA-GA

GCAACGCGAAGAACCTTACC|АААТСТТGACATCCTTTG-АСААСТСТАGAGATAGAGCCTTCCCCCTTCGGGACAA А|СТ-GA GCAACGCGAAGAACCTTACC|AGGCCTTGACATCCAATGAACT-TTCTAGAGATAGATTGGTGC-TTCGGGAACAT T GCAACGCGAAGAACCTTACC|TGGTCTTGACATCCACAGAACT-TTCCAGAGATGGATTGGTGCCTTCGGGAACTG T|GA-GA GCAACGCGAAGAACCTTACC|TGGTCTTGACATCCACAGAACT−TTCCAGAGAGTGGATTGGTGC−TTCGGGAACTG Т|GA−GA GCAACGCGAAGAACCTTACC|ТААGCTTGACATCCCAATGACATCTCCTTAATCGGA--GATCTTCTCGGGGACAT Т|GGTGA 



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#### Claims for the following Contracting States: AT, BE, CH, DE, DK, FR, GB, GR, IT, LI, LU, NL, SE

- 1. A method for generating a DNA probe specific for an organism to be identified and capable of distinguishing between genera and species in a non-empirical manner, said method comprising the steps of:
  - a) amplifying, using a pair of oligonucleotide primers, a variable intergenic region of the genome present in the organism to be identified and in a number of organisms phylogenetically related to, or suspected of being present in a given sample containing said organism to be identified and having said varaible region in its genome, said printers corresponding to regions or portions thereof flanking said intergenic region and known or suspented of being conserved in said organisms;
  - b) determining the sequence of the amplified region;
  - c) selecting said sequence or a portion thereof to generate said probe specific for said organism to be identified by comparison with said other amplified regions; and
  - d) defining the hybridization conditions required to obtain a specific signal based on the precise nucleotide sequence of the selected probe.
- A method according to Claim 1, wherein one of said primers corresponds to a conserved region of the 16S rRNA gene and the other said primer corresponds to a conserved region of the 23S rRNA gene.
  - 3. A method according to Claim 1, wherein the primer sequences are derived from the ribonuclease P component RNA.
  - 4. A method according to any preceding claim, wherein the variable region amplified is derived from an autoclaved sample containing the organism to be identified.
- A method according to any preceding claim, wherein the microorganism is selected from a species of <u>Aeromonas</u>,
   Bacillus, Clostridium, Enterococcus, Eschericia, Klebsiella, Mycobacterium, Pseudomonas, Salmonella Serratia,
   Staphylococcus or <u>Streptococcus</u>.
  - 6. A DNA probe for <u>Clostridium difficile</u> obtained from a variable intergenic region intermediate the genes coding for 16S rRNA and 23S rRNA and derived from the following nucleotide sequence or a sequence complementary thereto:

SEQ ID NO: 1

- 7. A DNA probe for <u>Clostridium pasteurianum</u> obtained from a variable intergenic region intermediate the genes coding for 16S rRNA and 23S rRNA and derived from the following nucleotide sequence or a sequence complementary thereto:

SEQ ID NO: 2

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8. A DNA probe for <u>Clostridium perfringens</u> obtained from a variable intergenic region intermediate the genes coding for 16S rRNA and 23S rRNA and derived from the following nucleotide sequence or a sequence complementary thereto:

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. SEQ ID NO: 3

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 A DNA probe for <u>Mycobacterium avium</u> obtained from a variable intergenic region intermediate the genes coding for 16S rRNA and 23S rRNA and derived from the following nucleotide sequence or a sequence complementary thereto:

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SEQ ID NO: 4

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10. A DNA probe for <u>Mycobacterium bovis</u> obtained from a variable intergenic region intermediate the genes coding for 16S rRNA and 23S rRNA and derived from the following nucleotide sequence or a sequence complementary thereto:

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SEQ ID NO: 5

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11. A DNA probe for Mycobacterium tuberculosis obtained from a variable intergenic region intermediate the genes

coding for 16S rRNA and 23S rRNA and derived from the following nucleotide sequence or a sequence complementary thereto:

#### 5 SEQ ID NO: 6

ACCGGAAGTCGTCGGGATCACCTCCTTTCTAAGGAGCACCACGAAAACGCCCCAACTGG
TGGGTCAGGCGTGAGGGGTTCTTGTCTGTAGTGGGCGAGACGGGGTGCATGACAACAAA
GTTGGCCACCAACACACTGTTGGATCCTGAGGCAACACTCGGACTTGTTCCAGGTGTTG
TCCCACCGCCTTGGTGGTGGTGTGTGTTTTGAGAACGTGATAGTGGTTGCGAGCATCA
ATGGATACCCGTGCCGGCTAGCGGTGCCGTGTTCTTTGTGCAATATCTTTGGTTTTTGT
TGTGTTTGTAAGTGTCTAAGGGCGCA

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- 12. A method for the detection and/or determination of an organism containing or suspected of containing a target nucleotide sequence, which method comprises contacting said organism with a probe complementary to said target sequence according to any one of Claims 6-11.
- 20 13. A method of detecting a specific organism in a mixture of organisms, which method comprises determining the sequences of two selected nucleotide regions of said specific organism by a sequence of steps according to any one of Claims 1-5, and using said sequenced regions to generate a pair of primers to selectively amplify a specific nucleotide region of said organism and thereby detect said specific organism.

#### Claims for the following Contracting State: ES

- A method for generating a DNA probe specific for an organism to be identified and capable of distinguishing between genera and species in a non-empirical manner, said method comprising the steps of:
  - a) amplifying, using a pair of oligonucleotide primers, a variable intergenic region of the genome present in the organism to be identified and in a number of organisms phylogenetically related to, or suspected of being present in a given sample containing said organism to be identified and having said variable region in its genome, of said primers corresponding to regions or portions thereof flanking said intergenic region and known or suspected of being conserved in said organisms;
  - b) determining the sequence of the amplified region;
  - c) selecting said sequence or a portion thereof to generate said probe specific for said organism to be identified by comparison with said other amplified regions; and
  - d) defining the hybridization conditions required to obtain a specific signal based on the precise nucleotide sequence of the selected probe.
- A method according to Claim 1, wherein one of said primers corresponds to a conserved region of the 16S rRNA gene and the other said primer corresponds to a conserved region of the 23S rRNA gene.
- 45 3. A method according to Claim 1, wherein the primer sequences are derived from the ribonuclease P component RNA.
  - A method according to any preceding claim, wherein the variable region amplified is derived from an autoclaved sample containing the organism to be identified.

- A method according to any preceding claim, wherein the microorganism is selected from a species of <u>Aeromonas</u>, <u>Bacillus</u>, <u>Clostridium</u>, <u>Enterococcus</u>, <u>Eschericia</u>, <u>Klebsiella</u>, <u>Mycobacterium</u>, <u>Pseudomonas</u>, <u>Salmonella Serratia</u>, <u>Staphylococcus</u> or <u>Streptococcus</u>.
- 6. A method according to claim 1, wherein a DNA probe for <u>Clostridium difficile</u> is obtained from a variable intergenic region intermediate the genes coding for 16S rRNA and 23S rRNA and is derived from the following nucleotide sequence or a sequence complementary thereto:

SEQ ID NO: 1

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7. A method according to claim 1, wherein a DNA probe for <u>Clostridium pasteurianum</u> is obtained from a variable intergenic region intermediate the genes coding for 16S rRNA and 23S rRNA and is derived from the following nucleotide sequence or a sequence complementary thereto:

SEQ ID NO: 2

8. A method according to claim 1, wherein a DNA probe for <u>Clostridium perfringens</u> is obtained from a variable intergenic region intermediate the genes coding for 16S rRNA and 23S rRNA and is derived from the following nucleotide sequence or a sequence complementary thereto:

#### SEQ ID NO: 3

9. A method according to claim 1, wherein a DNA probe for Mycobacterium avium is obtained from a variable intergenic region intermediate the genes coding for 16S rRNA and 23S rRNA and is derived from the following nucleotide sequence or a sequence complementary thereto:

#### SEQ ID NO: 4

10. A method according to claim 1, wherein a DNA probe for <u>Mycobacterium bovis</u> is obtained from a variable intergenic region intermediate the genes coding for 16S rRNA and 23S rRNA and is derived from the following nucleotide sequence or a sequence complementary thereto:

11. A method according to claim 1, wherein a DNA probe for <u>Mycobacterium tuberculosis</u> is obtained from a variable intergenic region intermediate the genes coding for 16S rRNA and 23S rRNA and is derived from the following nucleotide sequence or a sequence complementary thereto:

#### SEQ ID NO: 6

- ACCGGAAGTCGTCGGGATCACCTCCTTTCTAAGGAGCACCACGAAAACGCCCCAACTGG
  TGGGTCAGGCGTGAGGGGTTCTTGTCTGTAGTGGGCGAGACGGGGTGCATGACAACAAA
  GTTGGCCACCAACACACTGTTGGATCCTGAGGCAACACTCGGACTTGTTCCAGGTGTTG
  TCCCACCGCCTTGGTGGTGGTGTGTGTTTTGAGAACGTGATAGTGGTTGCGAGCATCA
  ATGGATACCCGTGCCGGCTAGCGGTGGCGTGTTCTTTGTGCAATATCTTTGGTTTTTGT
  TGTGTTTTGTAAGTGTCTAAGGGCGCA
- 12. A method for the detection and/or determination of an organism containing or suspected of containing a target nucleotide sequence, which method comprises contacting said organism with a probe complementary to said target sequence according to any one of Claims 6-11.
  - 13. A method of detecting a specific organism in a mixture of organisms, which method comprises determining the sequences of two selected nucleotide regions of said specific organism by a sequence of steps according to any one of Claims 1-5, and using said sequenced regions to generate a pair of primers to selectively amplify a specific nucleotide region of said organism and thereby detect said specific organism.

#### 45 Patentansprüche

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## Patentansprüche für folgende Vertragsstaaten : AT, BE, CH, DE, DK, FR, GB, GR, IT, LI, LU, NL, SE

- Verfahren zur Erzeugung einer DNA-Sonde, welche für einen zu identifizierenden Organismus spezifisch ist und zwischen Gattungen und Arten auf eine nicht-empirische Weise unterscheiden kann, wobei das Verfahren die folgenden Schritte umfaßt:
- a) Amplifizierung eines variablen intergenischen Bereichs des Genoms, welcher in dem zu identifizierenden Organismus und in einer Anzahl von phylogenetisch verwandten Organismen vorhanden ist oder von dem vermutet wird, daß er in einer bestimmten Probe vorliegt, welche den zu identifizierenden Organismus enthält, der diesen variablen Bereich in seinem Genom aufweist, unter Verwendung eines Oligonukleotid-Primerpaares, wobei die Primer den intergenischen Bereich flankierenden Bereichen oder Teilen davon entsprechen,

von denen bekannt ist oder von denen vermutet wird, daß sie in den Organismen konserviert sind; b) Bestimmen der Sequenz des amplifizierten Bereichs;

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- c) Auswählen der Sequenz oder eines Teils davon zur Erzeugung der Sonde, welche für den zu identifizierenden Organismus spezifisch ist, durch das Vergleichen mit den anderen amplifizierten Bereichen: und
   d) Definieren der zur Erzeugung eines spezifischen Signals erforderlichen Hybridisierungsbedingungen unter Zugrundelegung der genauen Nukleotidsequenz der ausgewählten Sonde.
- 2. Verfahren nach Anspruch 1, wobei einer der Primer einem konservierten Bereich des 16S-rRNA-Gens entspricht und der andere Primer einem konservierten Bereich des 23S-rRNA-Gens entspricht.
- 3. Verfahren nach Anspruch 1, wobei die Primer-Sequenzen von der RNA-Ribonuclease P-Komponente abgeleitet sind.
- 4. Verfahren nach einem der vorstehenden Ansprüche, wobei der amplifizierte variable Bereich von einer autoklavierten Probe abgeleitet wird, welche den zu identifizierenden Organismus enthält.
  - Verfahren nach einem der vorstehenden Ansprüche, wobei der Mikroorganismus aus einer Art von Aeromonas, Bacillus, Clostridium, Enterococcus, Escherichia, Klebsiella, Mycobacterium, Pseudomonas, Salmonella, Serratia, Staphylococcus oder Streptococcus ausgewählt ist.
  - 6. DNA-Sonde gegen Clostridium difficile, welche aus einem Zwischenprodukt des variablen intergenischen Bereichs der die 16S-rRNA und 23S-rRNA codierenden Gene hergestellt und von der folgenden Nukleotidsequenz oder einer dazu komplementären Sequenz abgeleitet wurde:

- 7. DNA-Sonde gegen <u>Clostridium PAsteurianum</u>, welche aus einem Zwischenprodukt des variablen intergenischen Bereichs der die 16S-rRNA und 23S-rRNA codierenden Gene hergestellt und von der folgenden Nukleotidsequenz oder einer dazu komplementären Sequenz abgeleitet wurde:
- SEQ ID NO: 2

  AGAGAACCTGCCGGCTGGATCACCTCCTTTCTAAGGAGTAATTGTAGCAGGATAACTGT

  TGTATACATTGGTTTCTTACTCTTGTCTCTGTTTAATTTTGAGAGATCAGTTCTCTTAA

  GATGTACTTTGAAAATTGCATAGAGAAACAAAGTAAAGAAAAAATAATCCTTTGATAA

  TATGATTTTAATCGAAAAGATTGAAATTAAACAATAAAGACTAAAACTCTAAAACGGGCT

  AACGCCTAAAAGAGTAACAAGGTCAAGCTACAAAGGGCGCAT
  - 8. DNA-Sonde gegen Clostridium perfringens, welche aus einem Zwischenprodukt des variablen intergenischen Bereichs der die 16S-rRNA und 23S-rRNA codierenden Gene hergestellt und von der folgenden Nukleotidsequenz oder einer dazu komplementären Sequenz abgeleitet wurde:

SEQ ID NO: 3

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- 9. DNA-Sonde gegen Mycobacterium avium, welche aus einem Zwischenprodukt des variablen intergenischen Bereichs der die 16S-rRNA und 23S-rRNA codierenden Gene hergestellt und von der folgenden Nukleotidsequenz oder einer dazu komplementären Sequenz abgeleitet wurde:
- DNA-Sonde gegen <u>Mycobacterium bovis</u>, welche aus einem Zwischenprodukt des variablen intergenischen Bereichs der die 16S-rRNA und 23S-rRNA codierenden Gene hergestellt und von der folgenden Nukleotidsequenz oder einer dazu komplementären Sequenz abgeleitet wurde:
- 11. DNA-Sonde gegen Mycobacterium tuberculosis, welche aus einem Zwischenprodukt des variablen intergenischen Bereichs der die 16S-rRNA und 23S-rRNA codierenden Gene hergestellt und von der folgenden Nukleotidsequenz oder einer dazu komplementären Sequenz abgeleitet wurde:

SEQ ID NO: 6

ACCGGAAGTCGTCGGGATCACCTCCTTTCTAAGGAGCACCACGAAAACGCCCCAACTGG
TGGGTCAGGCGTGAGGGGTTCTTGTCTGTAGTGGGCGAGACGGGGTGCATGACAACAAA
GTTGGCCACCAACACACTGTTGGATCCTGAGGCAACACTCGGACTTGTTCCAGGTGTTG
TCCCACCGCCTTGGTGGTGGTGTGTTTTGAGAACGTGATAGTGGTTGCGAGCATCA
ATGGATACCCGTGCCGGCTAGCGGTGGCGTGTTCTTTGTGCAATATCTTTGGTTTTTGT
TGTGTTTGTAAGTGTCTAAGGGCGCA

- 12. Verfahren zum Nachweis und/oder zur Bestimmung eines Organismus, welcher eine Ziel-Nukleotidsequenz enthält oder von dem vermutet wird, daß er sie enthält, wobei das Verfahren das Inkontaktbringen des Organismus mit einer Sonde nach einem der Ansprüche 6-11 umfaßt, die zu der Zielsequenz komplementär ist.
  - 13. Verfahren zum Nachweis eines spezifischen Organismus in einer Mischung von Organismen, wobei das Verfahren das Bestimmen der Sequenzen zweier ausgewählter Nukleotidbereiche des spezifischen Organismus durch eine Sequenz der Schritte nach einem der Ansprüche 1-5 und die Verwendung der sequenzierten Bereiche zur Erzeugung eines Primerpaares umfaßt, um einen spezifischen Nukleotidbereich des Organismus selektiv zu amplifizieren und um auf diese Weise den spezifischen Organismus nachzuweisen.

#### 25 Patentansprüche für folgenden Vertragsstaat: ES

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- Verfahren zur Erzeugung einer DNA-Sonde, welche für einen zu identifizierenden Organismus spezifisch ist und zwischen Gattungen und Arten auf eine nicht-empirische Weise unterscheiden kann, wobei das Verfahren die folgenden Schritte umfaßt:
  - a) Amplifizierung eines variablen intergenischen Bereichs des Genoms, welcher in dem zu identifizierenden Organismus und in einer Anzahl von phylogenetisch verwandten Organismen vorhanden ist oder von dem vermutet wird, daß er in einer bestimmten Probe vorliegt. welche den zu identifizierenden Organismus enthält, der diesen variablen Bereich in seinem Genom aufweist, unter Verwendung eines Oligonukleotid-Primerpaares, wobei die Primer den intergenischen Bereich flankierenden Bereichen oder Teilen davon entsprechen, von denen bekannt ist oder von denen vermutet wird, daß sie in den Organismen konserviert sind; b) Bestimmen der Sequenz des amplifizierten Bereichs;
  - c) Auswählen der Sequenz oder eines Teils davon zur Erzeugung der Sonde, welche für den zu identifizierenden Organismus spezifisch ist, durch das Vergleichen mit den anderen amplifizierten Bereichen; und d) Definieren der zur Erzeugung eines spezifischen Signals erforderlichen Hybridisierungsbedingungen unter Zugrundelegung der genauen Nukleotidsequenz der ausgewählten Sonde.
- 2. Verfahren nach Anspruch 1, wobei einer der Primer einem konservierten Bereich des 16S-rRNA-Gens entspricht und der andere Primer einem konservierten Bereich des 23S-rRNA-Gens entspricht.
- 3. Verfahren nach Anspruch 1, wobei die Primer-Sequenzen von der RNA-Ribonuclease P-Komponente abgeleitet sind
- 4. Verfahren nach einem der vorstehenden Ansprüche, wobei der amplifizierte variable Bereich von einer autoklavierten Probe abgeleitet wird, welche den zu identifizierenden Organismus enthält.
  - Verfahren nach einem der vorstehenden Ansprüche, wobei der Mikroorganismus aus einer Art von <u>Aeromonas</u>, <u>Bacillus</u>, <u>Clostridium</u>, <u>Enterococcus</u>, <u>Escherichia</u>, <u>Klebsiella</u>, <u>Mycobacterium</u>, <u>Pseudomonas</u>, <u>Salmonella</u>, <u>Serratia</u>, <u>Staphylococcus</u> oder <u>Streptococcus</u> ausgewählt ist.
  - 6. Verfahren nach Anspruch 1, wobei eine DNA-Sonde gegen <u>Clostridium difficile</u> aus einem Zwischenprodukt des variablen intergenischen Bereichs der die 16S-rRNA und 23S-rRNA codierenden Gene hergestellt und von der folgenden Nukleotidsequenz oder einer dazu komplementären Sequenz abgeleitet wird:

SEQ ID NO: 1

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7. Verfahren nach Anspruch 1, wobei eine DNA-Sonde gegen <u>Clostridium pasteurianum</u> aus einem Zwischenprodukt des variablen intergenischen Bereichs der die 16S-rRNA und 23S-rRNA codierenden Gene hergestellt und von der folgenden Nukleotidsequenz oder einer dazu komplementären Sequenz abgeleitet wird:

SEQ ID NO: 2

8. Verfahren nach Anspruch 1, wobei eine DNA-Sonde gegen <u>Clostridium perfringens</u> aus einem Zwischenprodukt des variablen intergenischen Bereichs der die 16S-rRNA und 23S-rRNA codierenden Gene hergestellt und von der folgenden Nukleotidsequenz oder einer dazu komplementären Sequenz abgeleitet wird:

SEQ ID NO: 3

9. Verfahren nach Anspruch 1, wobei eine DNA-Sonde gegen Mycobacterium avium aus einem Zwischenprodukt des variablen intergenischen Bereichs der die 16S-rRNA und 23S-rRNA codierenden Gene hergestellt und von der folgenden Nukleotidsequenz oder einer dazu komplementären Sequenz abgeleitet wird:

SEQ ID NO: 4

10. Verfahren nach Anspruch 1, wobei eine DNA-Sonde gegen Mycobacterium bovis aus einem Zwischenprodukt des

variablen intergenischen Bereichs der die 16S-rRNA und 23S-rRNA codierenden Gene hergestellt und von der folgenden Nukleotidsequenz oder einer dazu komplementären Sequenz abgeleitet wird:

SEQ ID NO: 5

ACCGGAAAGGTCCGTGCGTGAAATTTAACCTTCCTTCCCTTTCTAAGGAGCACCACGA
AAACGCCCCAACTGGTGGGCGTAGGCGTGAGGGGTTCTTGTCTGTAGTGGGCGAGACGG
GGTGCATGACAACAAAGTTGCCACCAACACACTGTTGGGTCCTGAGGCAACACTCGGAC
TTGTTCCAGGTGTTGTCCCCACCGCCTTGGTTGGTGGGGTGTGTTTTGAGAACTGG
ATAGTGGTTGCGAGCATCAATGGATACGCTGCCGGCTAGCGGTGGCGTGTTCTTTGTGC
AATATTCTTTGGTTTTTGTTGTTGTTTTGTAAGTGTCTAAAGGGCCGCAT

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11. Verfahren nach Anspruch 1, wobei eine DNA-Sonde gegen <u>Mycobacterium tuberculosis</u> aus einem Zwischenprodukt des variablen intergenischen Bereichs der die 16S-rRNA und 23S-rRNA codierenden Gene hergestellt und von der folgenden Nukleotidsequenz oder einer dazu komplementären Sequenz abgeleitet wird:

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SEQ ID NO: 6

ACCGGAAGTCGTCGGGATCACCTCCTTTCTAAGGAGCACCACGAAAACGCCCCAACTGG
TGGGTCAGGCGTGAGGGGTTCTTGTCTGTAGTGGGCGAGACGGGGTGCATGACAACAAA
GTTGGCCACCAACACACTGTTGGATCCTGAGGCAACACTCGGACTTGTTCCAGGTGTTG
TCCCACCGCCTTGGTGGTGGTGTGTTTTGAGAACGTGATAGTGGTTGCGAGCATCA
ATGGATACCCGTGCCGGCTAGCGGTGGCGTGTTCTTTTGTGCAATATCTTTGGTTTTTGT
TGTGTTTGTAAGTGTCTAAGGGCGCA

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- 12. Verfahren zum Nachweis und/oder zur Bestimmung eines Organismus, welcher eine Ziel-Nukleotidsequenz enthält oder von dem vermutet wird, daß er sie enthält, wobei das Verfahren das Inkontaktbringen des Organismus mit einer Sonde nach einem der Ansprüche 6-11 umfaßt, die zu der Zielsequenz komplementär ist.
- 13. Verfahren zum Nachweis eines spezifischen Organismus in einer Mischung von Organismen, wobei das Verfahren das Bestimmen der Sequenzen zweier ausgewählter Nukleotidbereiche des spezifischen Organismus durch eine Sequenz der Schritte nach einem der Ansprüche 1-5 und die Verwendung der sequenzierten Bereiche zur Erzeugung eines Primerpaares umfaßt, um einen spezifischen Nukleotidbereich des Organismus selektiv zu amplifizieren und um auf diese Weise den spezifischen Organismus nachzuweisen.

#### 45 Revendications

Revendications pour les Etats contractants sulvants : AT, BE, CH, DE, DK, FR, GB, GR, IT, LI, LU, NL, SE

- Méthode de génération d'une sonde d'ADN spécifique d'un organism à identifier et capable de distinguer entre les genres et les espèces de manière non-empirique, ladite méthode comprenant les étapes consistant à:
  - a) amplifier, en utilisant un couple d'amorces oligonucléotidiques, une région intergénique variable du génome présent dans l'organisme à identifier et dans un certain nombre d'organismes phylogénétiquement apparentés, ou susceptible d'être présent dans un échantillon donné contenant ledit organisme à identifier et ayant ladite région variable dans son génome, lesdites amorces correspondant à des régions ou à des parties de ces régions adjacentes à ladite région intergénique, connues ou susceptibles d'être conservées dans lesdits organismes;

- b) déterminer la séquence de la région amplifiée;
- c) sélectionner ladite séquence ou une partie de celle-ci pour générer ladite sonde spécifique dudit organisme à identifier par comparaison avec d'autres régions amplifiées ; et
- d) définir les conditions d'hybridation requises pour obtenir un signal spécifique basé sur la séquence nucléotidique précise de la sonde sélectionnée.
- Méthode selon la revendication 1, dans laquelle l'une desdites amorces correspond à une région conservée du gène de l'ARNr 16S et l'autre desdites amorces correspond à une région conservée du gène de l'ARNr 23S.
  - Méthode selon la revendication 1, dans laquelle les séquences d'amorce dérivent du composant ARN de la ribonucléaseP.
- 4. Méthode selon l'une quelconque des revendications précédentes, dans laquelle la région variable amplifiée dérive d'un échantillon qui a été traité en autoclave et contient l'organisme à identifier.
  - Méthode selon l'une quelconque des revendications précédentes, dans laquelle le micro-organisme est sélectionné parmi les espèces Aeromonas, Bacillus, Clostridium, Enterococcus, Escherichia, Klebsiella, Mycobacterium, Pseudomonas, Salmonella, Serratia, Staphylococcus ou Streptococcus.
    - 6. Sonde d'ADN pour Clostridinm difficile obtenue à partir d'une région intergénique variable intermédiaire entre les gènes codant pour l'ARNr 16S et l'ARNr 235 et dérivée de la séquence nucléotidique suivante ou d'une séquence complémentaire de celle-ci

#### SEQ ID NO: 1

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## GAATAAAGTAAAGCTAAAGGTATATAAAAATCCTTTGTAAGAATCAATTT AAGGTCAAGCTACAAAGGGCGCAT

7. Sonde d'ADN pour Clostridium pasteurianum obtenue à partir d'une région intergénique variable intermédiaire entre les gènes codant pour l'ARNr 16S et l'ARNr 23S et dérivée de la séquence nucléotidique suivante ou d'une séquence complémentaire de celle-ci

#### SEQ ID NO: 2

- - 8. Sonde d'ADN pour Clostridium perfringens obtenue à partir d'une région intergénique variable intermédiaire entre les gènes codant pour l'ARNr 16S et l'ARNr 235 et dérivée de la séquence nucléotidique suivante ou d'une séquence complémentaire de celle-ci

#### SEQ ID N° 3

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- 25 AGGAGAACTGCGGCTGGATCACCTCCTTTCTAAGGAATACATCTTAGGA
  CAACTAAGATGATAATGAATTCTGGATAATATCTCTGTTTAATTTTGAGA
  GACTATCTCTCAAAATTGTTCTTTGAAAATTGCACATAATTTAATTTATA
  GAAACAACAAGCCAAATTGGCAAAACCAATTTCTATTCTTTGTAAAATGA
  GAACTATAACTAATATAGGTCAAGCTACAAAGGGCCGCAT
- 9. Sonde d'ADN pour Mycobacterinm avium obtenue à partir d'une région intergénique variable intermédiaire entre les gènes codant pour l'ARNr 16S et l'ARNr 23S et dérivée de la séquence nucléotidique suivante ou d'une séquence complémentaire de celle-ci

## seq ID No: 4

- ACCGGAGGTGGGGCTGATCACCTCCTATTCTAAGAGCACCACAAAACGA

  CCCCGAACTGGTGGGGTCGGGAGCCAGTAGGGGTTCCCGTCTAGTGAC

  GGGGGCCGGGTGCGCAACAGAAATGATTGCCAGACACACTATTGGGCC

  CTGAGACAACACTCGGTCCGTCCGTGTGGAGTCCCTCCATCTTGGTGGT

  GGGGTGTGTGGTGTTTGTATTGGGATAGTGGTTGCGATGATCTAGGTGA

  GCGCATGGTCTTCGTGGCCGGCCGTTGATCGAAATGGGTAATTTCTTTT

  TTAACTCTTGTGTGTAAGTAAGTGTTTAAGGGGGGGAT
  - 10. Sonde d'ADN pour Mycobacterium bovis obtenue à partir d'une région intergénique variable intermédiaire entre les gènes codant pour l'ARNr 16S et l'ARNr 23S et dérivée de la séquence nucléotidique suivante ou d'une sé-



quence complémentaire de celle-ci

#### SEQ ID NO: 5

**ACCGGAAAGGTCCGTGCGTGAAATTTAACCTTCCTTCCCTTTTCTAAGG** AGCACCACGAAAACGCCCCAACTGGTGGGCGTAGGCGTGAGGGGTTCT TGTCTGTAGTGGGCGAGACGGGGTGCATGACAACAAGTTGCCACCAAC ACACTGTTGGGTCCTGAGGCAACACTCGGACTTGTTCCAGGTGTTGTCC CCACCGCCTTGGTTGGTGGGGTGTGTGTTTTGAGAACTGGATAGTGGT TGCGAGCATCAATGGATACGCTGCCGGCTAGCGGTGGCGTGTTCTTTGT GCAATATTCTTTGGTTTTTGTTGTTGTAAGTGTCTAAAGGGCGCAT

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11. Sonde d'ADN pour Mycobacterium tuberculosis obtenue à partir d'une région intergénique variable intermédiaire entre les gènes codant pour l'ARNr 16S et l'ARNr 23S et dérivée de la séquence nucléotidique suivante ou d'une séquence complémentaire de celle-ci

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#### SEQ ID NO: 6

ACCGGAAGTCGTCGGGATCACCTCCTTTCTAAGGAGCACCACGAAAACG 30 CCCCAACTGGTGGGTCAGGCGTGAGGGGTTCTTGTCTGTTAGTGGGCG **AGACGGGGTGCATGACAACAAGTTGGCCACCAACACACTGTTGGATCC** TGAGGCAACACTCGGACTTGTTCCAGGTGTTGTCCCACCGCCTTGGTGG 35 TGGGTGTGTGTTGAGAACGTGATAGTGGTTGCGAGCATCAATGGATA CCCGTGCCGGCTAGCGGTGGCGTGTTCTTTGTGCAATATCTTTGGTTTT 40 TGTTGTGTTGTAAGTGTCTAAGGGCGCA

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- 12. Méthode de détection et/ou de détermination d'un organisme contenant ou susceptible de contenir une séquence nucléotidique cible, ladite méthode comprenant la mise en contact dudit organisme avec une sonde complémentaire de ladite séquence cible selon l'une quelconque des revendications 6-11.
- 13. Méthode de détection d'un organisme particulier dans un mélange d'organismes, ladite méthode comprenant la détermination des séquences de deux régions nucléotidiques sélectionnées dudit organisme particulier par une suite d'étapes selon l'une quelconque des revendications 1-5 et l'utilisation desdites régions séquencées pour générer un couple d'amorces afin d'amplifier sélectivement une région nucléotidique spécifique dudit organisme et de détecter ainsi ledit organisme spécifique.

#### Revendications pour l'Etat contractant sulvant : ES

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Méthode de génération d'une sonde d'ADN spécifique d'un organisme à identifier et capable de distinguer entre les genres et les espèces de manière non-empirique, ladite méthode comprenant les étapes consistant à :

- a) amplifier, en utilisant un couple d'amorces oligonucléotidiques, une région intergénique variable du génome présent dans l'organisme à identifier et dans un certain nombre d'organismes phylogénétiquement apparentés, ou susceptible d'être présent dans un échantillon donné contenant ledit organisme à identifier et ayant ladite région variable dans son génome, lesdites amorces correspondant à des régions ou à des parties de ces régions adjacentes à ladite région intergénique, connues ou susceptibles d'être conservées dans lesdits organismes;
- b) déterminer la séquence de la région amplifiée;
- c) sélectionner ladite séquence ou une partie de celle-ci pour générer ladite sonde spécifique dudit organisme à identifier par comparaison avec d'autres régions amplifiées ; et
- d) définir les conditions d'hybridation requises pour obtenir un signal spécifique basé sur la séquence nucléotidique précise de la sonde sélectionnée.
- Méthode selon la revendication 1, dans laquelle l'une desdites amorces correspond à une région conservée du gène de l'ARNr 16S et l'autre desdites amorces correspond à une région conservée du gène de l'ARNr 23S.
- Méthode selon la revendication 1, dans laquelle les séquences d'amorce dérivent du composant ARN de la ribonucléaseP.
  - 4. Méthode selon l'une quelconque des revendications précédentes, dans laquelle la région variable amplifiée dérive d'un échantillon qui a été traité en autoclave et contient l'organisme à identifier.
- 5. Méthode selon l'une quelconque des revendications précédentes, dans laquelle le micro-organisme est sélectionné parmi les espèces Aeromonas, Bacillus, Clostridium, Enterococcus, Escherichia, Klebsiella, Mycobacterium, Pseudomonas, Salmonella, Serratia, Staphylococcus ou Streptococcus
- 6. Méthode selon la revendication 1, dans laquelle une sonde d'ADN pour Clostridium difficile est obtenue à partir d'une région intergénique variable intermédiaire entre les gènes codant pour l'ARNr 16S et l'ARNr 23S et est dérivée de la séquence nucléotidique suivante ou d'une séquence complémentaire de celle-ci :

#### SEQ ID NO: 1

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7. Méthode selon la revendication 1, dans laquelle une sonde d'ADN pour Clostridium pasteurianum est obtenue à partir d'une région intergénique variable intermédiaire entre les gènes codant pour l'ARNr 16S et l'ARNr 23S et est dérivée de la séquence nucléotidique suivante ou d'une séquence complémentaire de celle-ci :

#### SEQ ID NO: 2

- - 8. Méthode selon la revendication 1, dans laquelle une sonde d'ADN pour *Clostridium perfringens* est obtenue à partir d'une région intergénique variable intermédiaire entre les gènes codant pour l'ARNr 16S et l'ARNr 23S et est dérivée de la séquence nucléotidique suivante ou d'une séquence complémentaire de celle-ci :

#### SEO ID N° 3

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- 25 AGGAGAACTGCGGCTGGATCACCTCCTTTCTAAGGAATACATCTTAGGA
  CAACTAAGATGATAATGAATTCTGGATAATATCTCTGTTTAATTTTGAGA
  GACTATCTCTCAAAATTGTTCTTTGAAAATTGCACATAATTTAATTTATA

  GAAACAACAAGCCAAATTGGCAAAACCAATTTCTTTTGTAAAATGA
  GAACTATAACTAATATAGGTCAAGCTACAAAGGGCGCAT
- 9. Méthode selon la revendication 1, dans laquelle une sonde d'ADN pour Mycobacterium avium est obtenue à partir d'une région intergénique variable intermédiaire entre les gènes codant pour l'ARNr 16S et l'ARNr 235 et est dérivée de la séquence nucléotidique suivante ou d'une séquence complémentaire de celle-ci :

#### 40 SEQ ID NO: 4

- ACCGGAGGTGGGGCTGATCACCTCCTATTCTAAGAGCACCACAAAACGA
  CCCCGAACTGGTGGGGTCGGGAGCCAGTAGGGGTTCCCGTCTAGTGAC
  GGGGGCCGGGTGCGCAACAGAAATGATTGCCAGACACACTATTGGGCC
  CTGAGACAACACTCGGTCCGTCCGTGTGGAGTCCCTCCATCTTGGTGGT
  GGGGTGTGTGTGTTTGTATTGGGATAGTGGTTGCGATGATCTAGGTGA
  GCGCATGGTCTTCGTGGCCGGCCGTTGATCGAAATGGGTAATTTCTTTT
  TTAACTCTTGTGTGTAAGTAAGTGTTTAAGGGGGGGAT
  - 10. Méthode selon la revendication 1, dans laquelle une sonde d'ADN pour Mycobacterium bovis est obtenue à partir d'une région intergénique variable intermédiaire entre les gènes codant pour l'ARNr 16S et l'ARNr 235 et est dérivée de la séquence nucléotidique suivante ou d'une séquence complémentaire de celle-ci:

#### SEQ ID NO: 5

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- 11. Méthode selon la revendication 1, dans laquelle une sonde d'ADN pour *Mycobacterium tuberculosis* est obtenue à partir d'une région intergénique variable intermédiaire entre les gènes codant pour l'ARNr 16S et l'ARNr 235 et est dérivée de la séguence nucléotidique suivante ou d'une séquence complémentaire de celle-ci:

#### SEQ ID NO: 6

- 12. Méthode de détection et/ou de détermination d'un organisme contenant ou susceptible de contenir une séquènce nucléotidique cible, ladite méthode comprenant la mise en contact dudit organisme avec une sonde complémentaire de ladite séquence cible selon l'une quelconque des revendications 6-11.
  - 13. Méthode de détection d'un organisme spécifique dans un mélange d'organismes, ladite méthode comprenant la détermination des séquences de deux régions nucléotidiques sélectionnées dudit organisme specifique par une suite d'étapes selon l'une quelconque des revendications 1-5 et l'utilisation desdites régions séquencées pour générer un couple d'amorces afin d'amplifier sélectivement une région nucléotidique spécifique dudit organisme et de détecter ainsi ledit organisme spécifique.

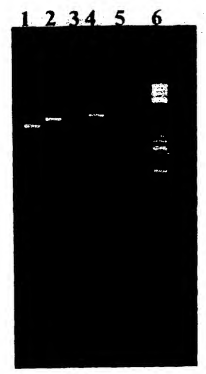


FIG.1

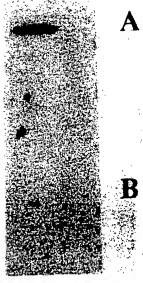


FIG.2

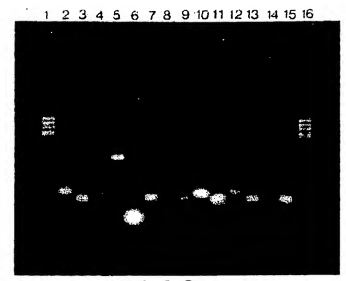


FIG.3

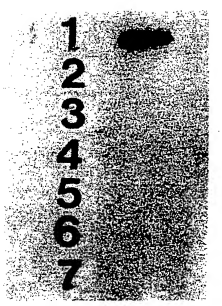


FIG.4

